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- (54) Substituted guinazoline derivatives
- (57) This invention provides a compound having the formula

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wherein:

X is phenyl which is optionally substituted;
R and R₁
Rependently, hydrogen, alleyl, allicoxy, hydroxy, or triftuoromethyt,
to hydrogen, alleyl, allicoxy, hydroxy, trifluoromethyt,

is a radical selected from the group consisting of

$$R_3 = R_3$$

H₃ is independently hydrogen, alkyl, carboxy, carboalkoxy, phenyl, or carboalkyl; n = 2-4;

or a pharmaceutically acceptable salt thereof, with the proviso that each R_3 of Y may be the same or different which are useful as antineoplestic agents.

Description

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This invention relates to cartain substituted quinazoline derivatives as well as the pharmacountically acceptable salts thereof. The compounds of the present invention inhibit the action of certain growth interest receptor profession tyrosine kinases (PTIQ) thereby inhibiting the abnormal growth of certain cell types. The compounds of this invention therefore are self-accors organise and are useful for the retainment of carpor in mammals. In addition, this invention relates to the manufacture of said quinazolines, their use for the freatment of cancer, and the pharmacountial preparations containing them.

Protein tyrosine kinases are a class of enzymes that catalyze the transfer of a phosphate group from ATP to a tyrosine residue located on a protein substrate. Protein tyrosine kinases clearly play a role in normal cell growth. Many of the growth factor receptor proteins function as tyrosine kinases and it is by this process that they effect signaling. The interaction of growth factors with these receptors is a necessary event in normal regulation of cell growth. However, under certain conditions, as a result of either mutation or overexpression, these receptors can become deregulated; the result of which is uncontrolled cell proliferation which can lead to lumor growth and ultimately to the disease known as cencer [Wilks A.F., Adv. Cancer Res., 60, 43 (1993) and Parsons, J.T.; Parsons, S.J., Important Advances in Oncology, DeVita V.T. Ed., J.B. Lippincott Co., Phila., 3 (1993) J. Among the growth factor receptor kinases and their protooncogeines that have been identified and which are targets of the compounds of this invention are the opidermal growth factor receptor kiriase (EGF-R kinase, the protein product of the erbB oncogene), and the product produced by the erbB-2 (also referred to as the neu or HER2) oncogene. Since the phosphorylation event is a necessary signal for cell division to occur and since overexpressed or mutated kinases have been associated with cancer, an inhibitor of this event, a protein tyrosine kinase inhibitor, will have therepeutic value for the treatment of cancer and other diseases. characterized by uncontrolled or abnormal cell growth. For example, overexpression of the receptor kinase product of the erbB-2 oncopene has been associated with human breast and ovarian cancers (Slamon, D. J., et. al., Science, 244, 707 (1969) and Science, 235, [145 (1987)]. Deregulation of EGF-FI kinase has been associated with epidermoid tumors [Reiss, M., et. al., Cancer Res., 51, 6254 (1991)], breast tumors [Maclas, A., et. al., Anticancer Res., 7, 459 (1987)], and tumors involving other major organs [Guillick, W.J., Brit. Med. Bull., 47, 87 (1991)]. Because the importance of the role played by deregulated receptor kinases in the pathogenesis of cancer, many recent studies have dealt with the development of specific PTK inhibitors as potential anti-cancer therapeutic agents Isome recent reviews: Burke. T.R., Drugs Future, 17, 119 (1992) and Chang, C.J.; Geahlen, R.L., J. Nat. Prod., 55, 1529 (1992)].

The compounds of this invention are certain 4-enitinoquinazolines. Throughout this patent application, the quinazoline ring system will be numbered as indicated in the formula below.

Other 4-antimoquinazolines which differ both in the nature and placement of the substituents at positions 5-6 compared to the compounds of this invention have been noted to have PTK inhibition activity. It is known from the European Patent Application 520,722 At certain 4-antilinoculnazolines which contain at positions 5-8 hydrogen, chloro, trifluoromethyl, or nitro substituents. None of the compounds in the aforementioned application have the unique combination of substituents contained in the compounds of the present invention, in addition, it is noteworthy that although an articancer utility is claimed for the compounds of the aforementioned European Patent Application, no demonstration of an in vivo anti-cancer effect is provided. It is known from the European Patent Application 566,226 A1 certain 4-anilinequinazolines which optionally contain at positions 5-8 a variety of substituents. None of the compounds in the aforementioned application have the unique combination of substituents contained in the compounds of the present invention, in addition, it is noteworthy that although an anti-cancer utility is claimed for the compounds of the aforementioned European Patent Application, no demonstration of an in vivo anti-cancer effect is provided. The only in vivo activity described in the aforementioned European Patent Application is the inhibition of TGF-alpha stimulated growth of hepatocyte in rats, It is known from the European Patent Application 635,498 A1 certain 4-anitinoquinazolines which optionally have at position 6 a variety of substituents while at position 7 they must have a halogen. None of the comcounds in the aforementioned application have the unique combination of substituents contained in the compounds of the present invention. In addition, it is noteworthy that although an anti-cancer utility is claimed for the compounds of the aforementioned European Petent Application, no demonstration of an in vivo anti-cancer effect is provided. The only in vivo activity described in the aforementioned European Patent Application is the inhibition of TGF-alpha stimulated growth of hepatocyte in rats. In addition, certain quinazoline inhibitors that do not have a 4-anilino group are known. It is known from the European Patent Application 602,851 A1 certain pulnazolines that do not have an apilline group in the 4 position and which potionally contain at positions 5-8 a variety of substituents. None of the compounds in the algrementioned application have the unique combination of substituents contained in the compounds of the present invention. In addition, it is noteworthy that although an anti-cancer utility is claimed for the compounds of the aforementioned European Patent Application, no demonstration of an in vivo anti-cancer effect is provided. The only in vivo activity described in the aforementioned European Patent Application is the inhibition of TGF-alpha stimulated growth of hepatocyte in rats. It is known from the patent application WO 95/19774 certain heterocycles that are inhibitors of PTKs that have a similar pyrimidine drig to the 4-anilinoquinazoles of the present invention. This aforementioned application makes no mention of 4-anifinoquinazolines nor of the unique combination of substituents contained in the compounds of the present invention. In addition, it is noteworthy that although an anti-cancer utility is claimed for the compounds of the aforementioned application, no demonstration of an in vivo anti-cancer effect is provided. It is known from the patent application WO 95/157581 certain quinazolines which optionally contain at positions 5-7 a variety of substituents. None of the compounds in the aforementioned application have the unique combination of substituents contained in the compounds of the present invention. In addition, it is noteworthy that although an artificancer utility is claimed for the compounds of the aforementioned patent application, no demonstration of an in vivo anti-cancer effect is provided.

In addition to the aloriementioning patient applications, a number of publications describe 4-sentinocquiracylines; Fry, D.W., et al., Science, 265, 1638 (1994), Flewcastle G.W., et. al., J. Med. Chem., 38, 3452 (1995), and Bridges, A.J., et. al., J. Med. Chem., 39, 627, [1995]). None of the compounds described in these publications have the unique combination of substituents contained in the compounds of the present (invention. In addition, it is noteworthy that no demonstration of an in vivo and scence effect is described in these proprist.

A PTK catalyses the transfer of a phosphale group from a molecule of ATP to a tyrosine residue located on a protein substrate. The inhibitors so far known in the art are usually competitive with either the ATP or the protein substrate of the kinase. Some of these inhibitors, the so-called mixed competitive inhibitors, can be competitive with both ATP and substrate simulteneously; all such competitive inhibitors function as reversible inhibitors. The 4-anilinoouinazolines known in the art are reversible inhibitors that are competitive with ATP (Fry, D.W., et. al., Science, 265, 1093 (1994)]. Since the concentration of ATP in a cell is normally very high (millimolar), compounds that are competitive with ATP may lack in vive activity since it is unlikely that said compounds can reach the concentrations within the cell that are necessary to displace the ATP from its binding site. As demonstrated, the quinazeline inhibitors of this invention have the unique ability of inhibiting these PTKs in an irreversible mannet and are therefore non-compelitive with ATP or protein substrate. The compounds of this invention can function as irreversible inhibitors by virtue of the fact that they can form covalent bonds to amino acid residues located at the active site of the enzyme. As shown below, this results in an enhanced therapeutic usefulness of the compounds of this invention when compared to the reversible type of inhibitor. In particular, it is shown that it is the unique nature and combination of substituents contained in the compounds of the present invention that lead to the irreversible binding of the inhibitor to the enzyme. These unique properties of the compounds of this invention contribute to their ability to inhibit the growth of human tumors in an in vivo model of cancer.

This invention provides a compound of formula 1;

wherein:

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X is phanyl optionally substituted with one or more substituents selected from the group consisting of halogen, alkyl of 1-6 carbon atoms, alloxy of 1-6 carbon atoms; or 1-6 carbon atoms;

R and R₁ are each, independently, hydrogen, halogen, alkyl of 1-6 carbon atoms, alkoxy of 1-6 carbon atoms, hydroxy, or trifluoromethyl;

H₂ is hydrogen, alkyl of 1-6 carbon atoms, alkoxy of 1-6 carbon atoms, hydroxy, trifluoromethyl;

Y is a radical selected from the following:

A₃ is independently hydrogen, elkyl of 1-5 carbon atoms; carboxy, carboalkoxy of 2-7 carbon atoms, phenyl, or carboalkyl of 2-7 carbon atoms;

n = 2-4;

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or a pharmaceutically acceptable salt thereof, with the proviso that each R₂ of Y may be the same or different.

The pharmaceutically acceptable satis are those derived from such organic and inorganic acids as acetic, texts, ditric, texteric, succinic, materic, materia, gluconic, hydrochloric, hydrobromic, phosphoric, nitrie, sulfuric, methanissulfonic, and similarly known acceptable acids.

The sityl portion of the alloys, althou, carinositicary, carbositivs, and alteracylswinno substituents include both straight thair as well as branched carbon chaha. Carbony las defined as a -CO₂H radical, Carbonitivs of 2.7 carbon atoms is defined as a -CO₂H radical, where R*1s an alkyl radical of 1-6 carbon atoms. Carbonalay is defined as a -COP* radical, where R*1s an alkyl radical of 1-6 carbon atoms. It is protected that it is ribbon. I do not be a carbon atoms in the carbonalay is defined as a -COP* radical, where R*1s are alkyl radical of 1-6 carbon atoms. When X is substituted, it is preferred that it is ribbon. I do not have a compound of this kinemation carbana as assymmetric center, this invention covers the individual R and S entantiomers as well as the racernate with respect to such compound.

Of the compounds of this invention, preferred members include those in which R, R³, and R² are hydrogen; and those in which R, R¹, and R² are hydrogen and X is either unsubstituted or monosubstituted with halogen or alkyl of 1.8 action atoms.

The preparation of the compounds of this invention encompassed by Formula 9 is described below in Flowsheet A where R_i , R_{p_i} , R_{p_i} , R_{p_i} , R_{p_i} , R_{p_i} , R_{p_i} and R_{p_i} are defined and R_{p_i} is alkyl of 1-6 carbon atoms (preferably isobutyl). Y' is a radical selected from the group consisting of:

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wherein each R's is independently hydrogen, alkyl of 1-6 carbon atoms, carboxy, carboalkoxy of 1-6 carbon atoms, phenyl, or carboalkyl of 2-7 carbon atoms. According to the sequence of reaction outlined in flowsheet A, a 5-offroanthranilonitrile of Formula 2 is heated at about 100°C with or without solvent containing an excess of dimethyllormamide dimethyl acetal to furnish an amidine of Formula 3. Heating a solution of amidine 3 and the antiline 4 in acetic acid for 1 to 5 hours gives the 6-nitro-4-anilinoquinazolines of Formula 5. Reduction of the nitro group of 5 using a reducing agent such as iron in an acctic acid-alcohol mixture at elevated temperature gives the 6-amino-4-anillinoquinazolines of Formula 6. Acylation of 6 with either an acid chloride of Formula 7 or a mixed anhydride of Formula 8 (which is prepared from the corresponding carboxylic acid) in an inert solvent such as tetrahydrofuran (THF) in the presence of an organic base such as pyridine or triethylamine gives the compounds of this invention represented by Formula 9, in those cases where 7 or 8 have an asymmetric carbon atom, they can be used as the recemate or as the individual R or Sentantiomers in which case the compounds of this invention will be in the racemic or R and S optically active forms, respectively. The 5-mitro-anthrepilionitriles of Formula 2 needed to prepare the compounds of this invention are either already known to the art or can be prepared by procedures known in the art as detailed in the following references; Baudet, Recl.Trav.Chim.Pays-Bas, 43, 710 (1924); Harlmans, Recl.Trav.Chim.Pays-Bas, 65, 468, 469 (1946); Taylor et al., J.Amer.Chem.Soc., 82, 6058,6063 (1960); Taylor et al., J.Amer.Chem.Soc., 82, 3152,3154 (1960); Deshpando; Seshadri, Indian J.Chem., 11, 538 (1973); Katritzky, Alan R.; Laurenzo, Kathleen S., J.Org.Chem., 51 (1986); Niclas, Hans-Joachim; Bohle, Matthias; Flick, Jens-Deltey, Zeuner, Frank; Zoelch, Lothar, Z.Chem., 25(4). 137-138 (1985).

FLOWSHEET A

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The preparation of the compounds of this invention encompassed by Formula 12 is described below in Firewsheet B wherein R, R₁, R₂, X and n are described above. Each R₃ is independently hydrogen, phenyl, or allyl of 1-6 extroations. According to the reaction outlined in Flowsheet B, the 6-amino-4-anifiroquirazolines of Formula 10 (prepared as in Flowsheet A) are explated with a cyclic enhyldride of Formula 11 in an inert solvent such as tetrahydiculuran in the presentice of a basic catalays such as pyrifidor or infethylamins.

FLOWSHEET B

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Representative compounds of this invention were evaluated in several slandard phermiscological test procedures that showed that the compounds of this invention possess significant activity as inhibitors of protein tyrosine kirases, and are antiprofiferative agents. Based on the activity shown in the standard phermiscological test procedures, the compounds of this invention are therefore discribed as antineoplastic agents. The test procedures used and results obtained are shown below.

The proparation of the compounds of this invanion encompassed by Formula 19 is described below in Flowshoet C A-choice 5-elimiquini-zolini, 13, (Morley, US, and Sinjesca, U. Chain. Soc., 950 (1949)) is noticed to 6-amino-4-chloropiulinazoline, 14, using a reducing again such as softurn hydrousilise in a two phases system consisting of tetrahydriculoran and water in repeasance of a small amount of phase transfer catalysts. Adjustion of 14 while other an acid chiedded Formula 15 or a nixed antividide of Formula 16 (which is propared from the corresponding carboxylis acid) in an inter solvent such as tetrahydricular or 14 may 16 may 1

FLOWSHEET C

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The preparation of the compounds of this invention encompassed by Formula 26 is described below in Flowshest. D wherein Y; R_b, and X are described above, According to the reactions coullined in Flowshest D, the attitute group of 20 (prepared as in Flewshest A) is reduced to the corresponding infinio compound 21 duling a palledium catalyst and a source of frydrogen which can be hydrogen itself or cyclohexens. Acytation of 21 with other an exist analysis of Formula 22 or a mixed analysistic of Formula 23 (which is prepared from the corresponding carboxylide acidy) and intervalvent such as tetrahydrofuran (THF) in the presence of an organic base such as pyridina or N-mothly indipholine gives the compounds of Formula 24 in those cases where 22 or 23 have an asymmetric carbon atom, they can be used as the readmentation as the individual For 5 entantiomers in which case the compounds of this invention will be in the recentic or R and 5 optically active forms, respectively. Heating a compound of Formula 24 with an antiline of Formula 25 in a fine stoyen such as arctic calci gives the compounds of this invention represented by Formula 25.

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FLOWSHEET D

Inhibition of Epidermal Growth Factor Receptor Kinase (EGF-R)

Test compounds were evaluated for their ability to inhibit the phosphorylation of the tyracine residue of a peptide auditation classifyed by the enzyme opidiarinal growth faster receptor tense. The peptide substante (FRI-SEG) has the sequence arg-arg-lot-die-giu-asp-tala-giu-tyr-ela-aria-arg-giy. This enzymo was obtained as a membrane extract (a A43) cells (Annotian Type Callure Collection, Flockville, MD). A431 cells were grown in 1716 flasts to 60% combinency. The cells were washed (vide-with prophasite buffered saline (PSS) without CS2-, Flasts were protect for 1.5 hours in 20 min PES with 1.0 mM entry-entrale ratio of the properties and contribuged at 000g for 10 minutes. The cells were substituted in 1 mil per 5 x 10° colls of cold lysis buffer (10mM 4-Cl-yydrocycythy)-1-pipera-insenteneautionic acid (HEPES), plf 1.6, 10 mM NBC1, 2mM EDTA, 1 mM phenytmethysulfouryl-flootside (PMSF), 10 mM pmfel aprolinn), 10 mg/ml elaporation, 0.1 mM sodium ontohovarisation in a Durine homogenizary with 10 stanks son ice, The yeare was centrifuged at 600g for 10 minutes first to clear cell debric and the supermatent further centrifuged at 000,000 g for 30 min at 4°C. The membrane pellet was suspended in 1.5 ml HMS Duffer (50 mM HEPES, plf 7.6, 12° mM MBC1, 10% giverol). The membrane pellet was subspracted in 1.5 ml HMS Duffer (50 mM HEPES, plf 7.6, 12° mM MBC1, 10% giverol). The membrane pellet was divided into allquots, immediately frozen in liquid nitrogen and attend at 70°C.

Test compounds were made into 10 mg/ml stock solutions in 100% dimethylsultoxide (DMSO). Prior to experiment, stock solutions were diluted to 500 mM with buffer (30 mM Hepes pH 7.4) and then serially diluted to the desired concentration.

An aliquet of the A431 membrane extract (10 mg/ml) was diluted in 30 mM HEPES (pH 7.4) to give a protein concentration of 50 u/ml. 10 4 pul of excipting preparallor, ESF (11 at 12 µg/ml) was added and incubated for 10 min on its followed by 4 µl of the test compound or buffer, this mix was incubated on itse for 50 min. To this was added has 39P-A7F (10 mCPml) diluted in 110 in assay buffer along with the substrate perigited at a concentration of 0.5 mM (control reactions got no test compound) and the reaction was allowed to proceed for 50 min at 30°C. The reaction was stopped with 10% TCA and foll on to lef for all least 10 min after which tubes were microcentifulged at full speed for 15 mis. The a portion of the supernatants were spotted on PB1 phosphocollulose discs and washed twice in 1% seetic add then water for 5 min such followed by scintillation counting. The inhibition data for representative compounds of the levention are shown below in TABLE 1. The IC₅₀ is the concentration of test compound needed to reduce the total around of phosphorpheted substraction by 50%. The % inhibition of the test compound was determined for at least three different concentrations and the IC₅₀ value was evaluated from the dose response curve. The % inhibition was evaluated with the following formula:

% inhibition = 100 - [CPM(drug)/CPM(control)] x 100

where CPM(drug) is in units of counts per minute and is a number expressing the amount of radiolabled ATP (§ 30P) incorporated dnot be R-PSRC peptide substitate by the exyment after 30 minutes at 30°C in the presence of test compound as measured by liquid scintillation counting. CPM(control) is in units of counts per minute and was a number expressing the amount of radiolabled ATP (a 3PP) incorporated onto the RR-SRC peptide substitate by the encytical 30 minutes at 30°C in the extension of test compound as measured by liquid seinfiltation counting. The GPM values were corrected for the background counts produced by ATP in the absence of the enzymatic reaction. The ICQ₀ values reported in TASIL It are averages of this number of tests condicided.

TABLE 1

Compound	IC50 (µM)	Number of Tests
Example 4	0.012	5
Example 5	0,198	4
Example 6	0.5	1
Example 7	0.05	t
Example 8	0.04	2
Example 9	0.002	20
Example 10	0.11	2
Example 11	0.056	4
Example 13	10-7	1
Example 14	10	1

Determination of Govalent Binding of Test Compound to Epidermal Growth Factor Receptor Kinase

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An aliquot of the A431 anzyme extract (prépaired às described abové) was cilluted to 50-100 jig/ml with 30 mM A100 per la containing E40 at 12 jug/ml concentration en that under standard test conditions approximately 2% reaction will take place and the final E67 concentration is 24 jug/ml (ain in the standard dassity describe abové). This mixture was insubstated at least 10 minutes at 4°C before use. This enzyme preparation was used for the following dialwhis test procedures.

To 60 µcf the enzyme preparation was ackded 48 µl of test compound dissolved in 5% dimethybulloxite (DMSO) (rpial 48 µbf 5). MoSt of the bottoni). Test compound descentrabilism were chosen to be 20-100 lodd above the 1650 (rot pial 40 µbf 5). MoSt of the bottoni). The enzymo-inhibitor solution was incubated for 45-50 minutes at 4°C. Fer the undistyzed control test procedures, a 9 µl aliqued of the anzyme-inhibitor solution was evaluated under siendard protocols as descended above. For displies tost procedures, a 60 µl aliquot of the enzyme-inhibitor solution was placed in a wild of a Pierze Microdialyzer System 100 and dislyzed at 4°C versus 30 mM Hepse containing 1.25 µg/ml ESF for 24 hours with two changes of buffer (minimum 3 hours dislyzed store each change). The 6000 molecular weight outoff membranes were used. A 9 µl (at least duplicates) eliquot of the dislyzed solution was evaluated for activity the standard protocol as described above. Enzyme without added extra compound retains 50-00% of its initial activity after claylas. Dislyzed solutions of test compound without added enzyme are also evaluated to ensure the compounds are delyzable.

If the enzymatic activity is not recovered after dialysis, then it is determined that the feet compounds is binding contently (provessible shibition). If the enzymatic activity is singly recovered after dialysis, then it is determined that the test compound is binding non-covalently (reversible inhibition). Determinations of covalent binding can be expressed as the %-Recovered Activity which is calculated using the following formula which utilize the %-inhibition before and after dialysis:

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% Recovered Activity = [(% inhibition{pre dialysis}-% inhibition{post dialysis])/% inhibition {pre dialysis}] X 100

A value for the %- Receivered Activity close to 10% indicates non-covalent binding (reversible inhibition). A value for the %-Receivered Activity much below 10% indicates covalent hinding (intrevensible inhibition). The results obtained for the determinations of covalent binding to EGF-R kinsse for representative compounds of this invention are provised below in TABLE 2. For comparison purposes, TABLE 2.8 lac provised so brinding sits on the %-brumophology-6.7-dimethic covar-quinazofinamine. This quinazofine inhibitor has been identified as a potent inhibitor of EGF-R kinsse [Fry. D. W., et. al., Science, 265. 1093 [1994]. Preveasel EG. W., et. al., V. Med. Chinar, 39, 2882 (1995), and Bridges, AJ., et. al., v. AJ., et. al., v. AJ., et. al., v. AJ., et. al., et. al., et. al

TABLES

		IADLL Z	a militar		
	Determination of Covalent Bi	nding to Epiderm	al Grov	nh Facto	or Receptor Kinase
15	Compound	% Rec Activit	overed	ı	Determination
	Example 4	20	11	17	covalent (irreversible binding)
20	Example 9	9	4		covalent (irreversible binding)
	N-(-3-bromophenyl)-5,7-dimethoxy- 4-quinazolinamine	102	70	107	non-covalent (reversible binding)

The results in TABLE 2 show that the compounds of this invention inhibit GEF-FI kingse in an irreversible manner by forming a coverent plinage to an almino said residue located on the enzyme, in this respect, they are distinctly different from the usual 4-unilinoquinazoilnes such as N-Q-bromophenyl-6,7-dimethoxy-4-quinazoilnamine which binds in a reversible manner. As will be defineded below, this difference in blinding abilities between the compounds of this invention and the issual quinazoilnes inhibitors of the prior and leads to significantly improved biological activity and therefore generate rhangeautic usefulness.

Inhibition of Cell Growth as Measured by the Incorporation of [3H]-Thymidine

Representative compounds of this invention were ovaluated for their ability to inhibit the growth of the cell lines described below in wire. The inhibition is quantitated by measuring the decrease in the incorporation of radio-bateful thyritidine when the cells are grown in this presence of the inhibition. A431 and SIGBR cell lines are obtained from American Type Collines Collection, Pockville, MD, Nove-197 colls recordinated by manifesting (NH 377 embessible through with an activated rat Nev oncepee. NHEK cells are obtained from Clonditic (Sen Diego, CA). Cells were routinely grown to a fundified production in SK Co.), and if These cell lines, are despendent on growth science which are ligance to the receptor tyrosine kineses that are the targets of the compounds of this invention, and have the following observated

- A431: human epidermoid carcinoma cells overexpressing EGFA
- Neu-9T3: NIH 3T9 cells transfected with activated Neu oncogene
- NHEK: EGF dependent normal human epidermal keratinocytes SKBR3: Human breast cancer cells overexpressing ErbB2 gene

The cell lines were grown in appropriate media as described below:

- A431: Dulbecco's Modified Eagles Media, high glucose, BRIJ/Gibco (10% Fetal Bovine Serum (FBS), Glutamine, Penicillin-Streptomycin) Dulbecco, R., Freeman, G. Virology 8, 396 (1959).
- Neu-3T3: Dulbeccos Modified Eagles Media, high glucose (10% Felal Bovine Serum, Glutamine, Penicillin-Streptomycin)
- SKBR3: Roswell Park Memorial Institute 1640 W/GLU (10% FBS, GLU, PS) Moore, G. E., Gerner, R. E. and Franklin, H. A. A.M.A. 199, 516 (1967).
- NHEK: Keratinocyte Growth Media, Clonetics Boyce, S.T. and Ham, Fl. G. In Vitro 17, 299 (Abstract No. 159) (1981)

Cells were seeded at 10,000 cells/well in 96 well plates in complete media and allowed to grow to tog phase. At

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this stage the complete medie was replaced with media containing 0.5% FBS (for cells growing in 10% FBS) or media lacking epidermial growth factor (FBF) (for cells growing in serum fee media). Alter overanifight incubation in low serum (or EGF lacking) media, the compounds to be evaluated were edded and cells remained in the presence of compounds (or 45 to 72 hours). Media with lest compound was then removed and complete media was added back. The cells were slowed to grow for 18 hours. This is followed by incubation in PHIIhymidine II inclinit in serum/EGF media) for 4 hours. Cells were yeaded in 0.5 M NeOH for at least 50 mile 327°C and rediscentify washipt of 0.5 M NeOH for at least 50 mile 327°C and rediscentify washipt.

The cell growth inhibition data is provided below in TABLE 3. The $(C_{\rm S})$ is the concentration of test compound needed to reduce the amount of PHI thymdrine interoperation by $SO_{\rm S}$. The S-inhibition of the compound overbiated was determined for at least three different concentrations and the $(C_{\rm S})$ value evaluated from the dose response curve. The S-inhibition is evaluated with the following insmitted.

% inhibition =100- ICPM(drug)/CPM(control)] x 100

where CPM(drug) is in units of counts per minute and is a number expressing the amount of [Pt]) hymridine incorporated into the DNA when cells are grown in the presence of feet compound as measured by liquid exhillation counting, CPM (control) is in units of counts per minute and is a number expressing the amount of [Pt]) hymridine incorporated onlo the DNA when cells are grown in the absence of feet compound as measured by tiquid schillation counting.

		TABLE 3		
Inhibition of Ca	ll Growth as Me	asprement by the l	ncomperation of [3	-I]-Thymidine (IC ₅₀
Compound	A431 (µM)	SKBR3 (µМ)	NHEK (µM)	NEU-3T3 (μM)
Example 4	0.07	> 50.	0.1.7	> 50
Example 5	0.825	0.30	0.17	10
Example 6	27	> 50	4,5	> 50
Example 7	0.45	5,5	0.45	7.5
Example 8	0.22	7	0,5	0.9
Example 9	0.011	1.057	0.002	0.002
Example 10	60	> 50	> 50	15
Example 11	9.8	4	0.85	0.4

In Vivo Inhibition of the Growth of Human Epidermoid Tumors (A431)

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SALB/c num temale mice (Charles River, Wilmington, MA) were used in the in vivo standard pharmacological test procedures. Human epitarimola caracinoma cells A-431 (American Type Culture Collection, Ricciville, Maryland # CRL-155) were grown in vitro as described above. A unit of \$2 X if of cells were lipseled SC little mice. When tumors attained a mess of botween 100 and 150 mg, the mice were randomized into treatment groups (day zero). Mice were frainted if Pence a days either on days 1, 5, and 9 or or days 1 through 10 post sleping with doses of either 60, 40 or 20 mg/kg/dose of the compound to be evaluated in 0.2% Klucel. Control antinate received no drug. Tumor mass was determined every 7 days ((leginal) X width?³2) for 25 days post stanging. Relative tumor growth (Mean tumor mass on day 7, 14, 21, and 28 divided by the mean tumor mass on day xero) is determined for each treatment group. The %17 C (Tumor/Control) is determined by dividing the relative tumor growth of the testade group by the relative tumor growth of the placebo group and multiplying by 100. A compound is considered to be active if the %170 is found to be ≤42%. The highlightin results obtained for the compound of Example 2 are provided below in TABLE 4.

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TABLE 4

Dose (mg/kg/ dose) «IP	RTG ^b	%T/C°	RTGb	%T/C°	RTG⁵	%T/C°	HTG ^b	%T/C°	S/T ^d
	Day 7		Day 16	ļ	Day 21		Day 28		-
Control	3.68		7,91		11.41		15.04		10/10
* BO	0.71	18	0.91	11	1.07	9	1,96	9	5/5
* 40	1.48	40	2.23	28	3.05	27	4.04	27	5/5
* 20	1.72	47	2.69	34	4,33	38	6,18	41	5/5
20	0.75	26	1.01	13	1.25	#1	2.53	17	5/5

a) Drugs administered IP on days 1, 5, 8 or on days 1 through 10 **.

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Relative Turnor Growth = Mean Turnor Mess on Day 7, 14, 21, 28

Mean Turnor Mess on Day 0

%T/C = Relative Turnor Growth of Treated Group X 100
Relative Turnor Growth of Plecebo Group

it) S/T = No., Survivors/No. Treated on Day +26 post tumor sleging.

The ability of the sompound of Example 9 and N-(3-bromopheny)-5.7-dimethoxy-4-quinazoitrámine to inhibit fise growth of human epidermoid tumors (A431) in vivo-are compared below in TABLE s. N-(3-Bromopheny)-5,7-dimethoxy-4-quinazoidramine was chosen as the comparison compound since this quinazoine inhibitor as been identified at a potent inhibitor of EGF-1 kinase (Fry, D.W., et. at., Science, 265, 1093 (1994); Rewasste G.W., et. al., J. Med. Chom., 39, 3422 (1995); Bridgos, A.J., et. al., J. Med. Chom., 39, 3422 (1995); Bridgos, A.J., et. al., J. Med. Chom., 39, 257 (1996)] and is shortingassed in the European Palent Application 565(228 At.).

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TABLE 5

Compound	RTG ^b Day 7	%T/C°	PTG bDay 14	%T/C*	HTG ^b Day 21	%T/C°	PTG ^b Day 28	%T/C°	Ş/T ^d
* Control	3.18	*	5.65	-	7:79		10,3		10/1
Example 9	1.11	35	1.26	22	1.51	1.9	2.55	22	14/1
N- (3-Bromophenyi)- 6,7-dimethoxy- 4-quinazolinamine	3,03	95	6,58	116	10,5	128	14.47	140	15/15

b)

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Relative Tumor Growth ≈ Mean Tumor Mass on Day 7, 14, 21, 28

Mean Tumor Mass on Day 0

9)

%T/C = Relative Turner Growth of Treated Group X 100 Relative Turner Growth of Placebo group

d) S/T = No. Sundvers/No. Treated on Day +28 post lumor staging.

As shown in TABLES 4-5, the compounds of this invention inhibit the growth of human tumes in mammals and any principle useful as antinepolastic agents, in this respect, tipy are distinctly different from the usual 4-antilinoquinazolines such as N-(3-bromopheny)-6,7-dimethoxy-4-quinazolinamine which is devoid of antineoplastic activity.

The ability of the compound of Example 8 is initible the growth of human epidermolit furnors (A431) <u>in vice</u> was compared wigh two structurally similar compounds Ny4- ((3-methylphenyl)amino)-8-quinazolinyl-tratemes to as Comparator A) and Ny4-((3-bronophenyl)amino)-8-quinazolinyl-butanemide (reterred to as Comparator B) which are covered by European Palant Applications 635,488A1 and 556,286 A1, respectfully. The results of these comparisons are shown, in Tables 5 and 7.

TABLE 6

							s (A431) in:mi
Compound ^a	RTG ^b Day	%T/C	FITG b Day	%T/C°	P(TGbDay 21	%T/6º	S/T ^d
Control	5,52		11,63		7.79		10/10

* Indicate shalidari digilificance of p < 0,01.

a) Drugs admitistered IP, Conflori and the 60 migling deses were administered on days 1, 6, and 9; 20 migling discre were administered by drugs to the confloring deserving deser

Relative Tumor Growth = Mean Tumor Mass on Day 7, 14, 21
Mean Tumor Mass on Day 0

%T/C = Flelative Tumor Growth of Treated Group X 100

d) S/T = No. Survivors/No. Treated on Day +21 post tumor staging.

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TABLE 6 (continued)

A Comparison of the In Vivo Inhibition by the Compound of Example 9 and N-[4-[(3-Methylphenyl)amino]-6-quinazoliny()-7-fluoro-2-propenamide (Comparator A) of the Growth of Human Epidermoid Tumors (A431) in mice. BTG^bDay %T/G° S/Td Compound^e RTG^bDay %T/CF HTG bDay %T/C¢ 14 21 10/10 18* 2.50 21* 3,77 24* Example 9 1.25 (80 mg/kg) Comparator 3.39 61 5.60 48 7.68 10/10 A (80 mg/kg) Example 9 0.79 14" 1.39 12* 2.55 16* 10/10 (20 mg/kg) 4.82 87 7.36 63 9/10 Comparator 8.75 56 A (20 mg/kg)

"Indicates statistical significance of $p \! \prec \! 0.01$.

a) Drugs administered IP. Control and the 60 mg/kg doses were administered on days 1, 5, and 9:20 mg/kg doses were administered on days 1 through 10.

Relative Tumor Growth = Mean Tumor Mass on Day 7, 14, 21
Mean Tumor Mass on Day 0

c)

%T/C = Helative Turnor Growth of Treated Group X 100

d) S/T = No. Survivors/No. Treated on Day +21 post tumor staging.

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TABLE 7

A Comparisor b	of the In Vivo								uinazoliny
Compounda	PTG ^b Day	%T/C°	HTGb Day 14	%T/C°	RTG ⁵ Day 21	%T/C°	PTG ^b Day 28	%T/C°	S/T ^d
Control	3,56		5.55		5.85		7,63		10/0
Example 9 (60 mg/kg)	0:89	25*	1,50	27*	2.44	42*	3,45	45*	5/5
Comparator B (80 mg/kg)	9.37	95	5.43	9B	6.21	106	10.26	142	5/5
Comparator B (20 mg/kg)	2.90	81	4.19	75	5.62	96	8,04	105	5/5

*Indicates statistical significance of p < 0.01.

a) Druge administered IP, Control and Se 80 mg/kg dozes were administered on days 1, 5, and 9; 20 mg/kg doze was administered on days 1 through 10.

b)

Relative Tumor Growth = Mean Tumor Mass on Day 7, 14, 21, 28

Mean Tumor Mass on Day 0

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%T/C = Relative Turnor Growth of Treated Group X 100 Relative Turnor Growth of Placebo Group

d) S/T = No; Survivors/No. Treated on Day +28 post tumor staging.

The results obtained in Tables 6 and 7 show that the compound of Example 9, a representative compound of this invention, significantly (o < 0.01) inhibited human i spidermoid tumor growth juvo; The structurally closest compounds of European Patent applications 635,488A1 (Comparator A) and 566,226 A1 (Comparator B) were substantially less active than the compound of Example 9, and both failed to significantly reduce tumor growth at both dosset setted.

Based on the results obtained for representative compounds of this invention, the compounds of this invention are particularly useful in treating, inhibiting the growth of, or eradicating neopleans. In particular, the compounds of this invention are useful in treating, inhibiting the growth of, or eradicating neopleans the texture seems EGFR such as those of the breast, isdemy, bladder, mobility, and prophegus, stomach, colon, cowary, or the press. Isomorphisms that express EGFR such as those of the breast, isdemy, bladder, mobility, larging, aspecting to the property of the property of the press is the property of the press is the property of the press is the property of the press of the press of the press of the press of the pressure of t

The compounds of this invention may formulated near or may be combined with one or more pharmeceutically acceptable centries for administration. For example, solvenis, disunds and the like, and may be administered orally in such forms as tablets, capsules, dispersible powders, granules, or suspensions containing, for example, from about 0.05 to 5% of sugar, and exists containing, for example, from about 0.05 to 5% of sugar, and exists containing, for example, from about 0.05 to 5% of sugar, and exists containing, for example, from about 0.05 to 5% expension of a single containing from about 0.05 to 5% expending agent in an isototic medium. Such pharmaceutical preparations may contain, for example, from about 0.05 to 5% exbest 0.05 up of the settive ingredient in combination with the centre, more usually between exbout 5% and 65% by weight.

The effective diseage of active ingredient employed may very depending on the particular compound employed, the mode of administration and the severity of the condition being irealed. However, in general, settletelory results are obtained when the compounds of the invention are administered at a deity dosege of from about 0.5 to about 100 mg/kg of animal body weight, optionally given in divided doses two follow fines a day, or in setsialand aflates from rost large mammals the total daily dosege is from about 0.5 to 100 mg of the active compound in fittinate admixture with a solid or fluid pharmaceutically acceptable carrier. This desege regimen may be adjusted to provide the optimal therepetit mappose. For example, several divided doses may be administered daily or the dose may be proportionally reduced as indicated by the edispendies of the thempount estimates.

These active compounds may be administered orally as well as by intravenous, intramuscular, or subculaneous routes. Solid carriers include starch, lactose, dicalchim phosphate, microcrystalline cellulidae, sucress and skedin, while lighd carriers include startle water, polyostyneyne glyools, non-jenic ovardacties, and odble oils such as corp, peanul and assame oils, as are appearated to the nature of the active ingredient and the particular form of administration desired. Adjuvants customarily employed in the preparation of pharmaceutical compositions may be advirtalizedually included, such as favoring agents, polaring agents, priserving agents, and antioxidants, for example, vitamin 6, esco-

bic acid, BHT and BHA.
The prefered pharmacoultiesl compositions from the standpoint of asset preparation and administration are solid compositions, particularly tablets and bard-filled or liquid-filled capsules. Oral administration of the compounds is preferred.

In some cases it may be desirable to extrinister the compounce directly to the sirvays in the form of an aerosel. These active compounds may also be administered prenetration for interpretability for interpretability of results of the second section of the section of th

The pharmaceutical forms suitable for injectable use include sterille aqueous solutions or dispersions and sterille powders for the extemporaneous preparation of sterille injectable solutions or dispersions. In all cases, the form must be sterille and must be full of the oxident that easy synghapiting volfst. In must be stable under the ponditions of misurature and storage and must be preserved against the contaminating action of microorganisms such as bacteria and tungt. The carrier can be a solvent or dispersion medium containing, for example, water, ethanol, polyol (e.g., glycero), propyleng glycol and liquid polystylenge glycol), suitable mixtures thereof, and vegetable quite.

For the treatment of cancer, the compounds of this invention can be administered in combination with other artitumor substances or with radiation. These other substances or radiation treatments can be given at the same or at different times as the compounds of this invention. These combined therapies may effect synergy and result in improved efficacy. For example, the compounds of this invention can be used in combination with mitotic whithers such as taxed or virbilastics, allystuling agents used as a delicant or cytechposamillo, antimitationalise such as 5 fluoriounced or hydroxyurea, DNA intercalators such as a driamycin or bleomycin, topoisomerase inhibitors such as seloposide or camptehod, and affiliatelizations such as termoxible.

The preparation of representative examples of the compounds of this invention is described below.

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Example 1

N'-(2-Cyano-4-nitrophenyl)-N,N-dimethylformamidine

A 40.8 g portion of 5-nitro-enthrentilonitrile and 40 ml of N, N-dimethyllormemide dimethyl acetal were heated on a steam beth for 2 hours. The solvents were removed at reduced pressive and the residue was taken up in methylene chloride. After passing this solution through Magnesol the solvent was removed. After washing with ether 50.8 g of N-(2-ovano-4-nitrophemy)-N,N-dimethyllormemidities was obtained.

io Example 2

N-(3-Bromophenyl)-6-nitro-4-quinazolinamine

A solution of 23.74 m of 3-bronno salfine and 40.5 p Nr(2-cyano4-nitophenyl)-NN-dimethyllomamidine in 100 n/ol of glacial excelsulacid was estimated and heated in an oil balle in 14.97 for 15 shorinz on cooling. Illifation of the resoluting solid gives a quantitative yield of N-(3-bronnophenyl)-6-nitro-4-quinazolinamine; mp = 267-270°C; mass spectrum (n/ ex. 345.

Example 3

N-(3-Bromophenyl)-4,6-quinazolindiamine

A mixture of 9.5 g of N-(3-bromophanyl)-8-rithtro-4-quinazofinamine and 18.8 g of into powder in 150 ml of sthanol and 150 ml of gladal acetic acid was heated in an oil ball at 120°C for 2 hours. After filtration of the solid, solid sodium ciarbonate was added to the filtrate giving a solid. This was littered, and the solid was extracted with methanol. The extracts were treated with charcoal and evaporated to a solid. After washing the solid with either 27.5 g of N-(5-bromophanyl-4,5-cultragofination was obtained: mass speciment (m/d): 915.

Example 4

4-[[4-](9-Bromophenyl)aminol-6-quinazolinyl[amino -4-oxo-(Z)-2-tiulenoic acid

A 15 ml portion of pyridine was added to 1,5 g of N4;5-bromophenyl;4-6-quinezolindramhe end 0.5 g of maleic solychida. Nitre stuffing overnight, the polvents were remixed on the rectory expensation; The solid leves taken put in should 4.00 ml of hot ethanica and the insolutely material titeratic lagive 0.35 g of 4.[44]-6-bromophenyljaminoj-8-quinazolinylj isnibig-4-bro-0/2;2-bullenote stuff; mass spectorum (mile); M+1 413, 415.

Example 5

4-[[4-[(3-Bromophenyl)amino]-6-quinazolinyl[amino]-4-oxo-(E)-2-butenoic acid, ethyl ester

A adultion IN 1(3-bromephenyl)-4,6-quinazolindamina in 15 mil of pyrithin was cooled in an ice bath and a colution of 1.22 go of strip fummy choicinds in 10 ml of maltylinen chicrids was added dropwise. After airing 16 or 1.5 hours, the reaction was allowed to come to room temperature. The solvents were removed at rickined pressure and the recicies was treated with water. The not exict was filtered and extracted into hota cations. After littration of the indebulber most read to the solution state of the solution state of the solution state of the contraction of the solution state of the contraction of the solution state of the solution in the solution state of the solution

Example 6

N-[4-[(3-Bromophenyl)amino]-6-quinazolinyl-3-methyl-2-butenamide

A solution of 1.58 g of N-(3-hromopheny)-4,6-quinazolindiamino in 15 ml of pyridino was cooled in an ice bath and a solution of 0.67 mt of 3.9-dimethylacrylcyl chlorido in 7 ml of either was added dropwise. After stirring and cooling for 2 hours, the solvents were removed at reduced pressure. The residue was treated with water and the respitting solid was recrystallized from methyl cellusolve to give 0.97 g of N-(4-(3-hromophenyl)amino)-8-quinazolinyl)-3-methyl-2-butenamide; me 3 000. 3010, mass spotrum (rwls: 305, 308).

Example 7

N-4-[(3-Bromophenyl)amino-6-quinazolinyl-(E)-2-butenamide

A solution of 1,8 g of N-{3-bromophenyl}-4,6-quinazolindiamine in 15 mi of pyridine was cooled in an ice batch and a solution of 0,57 mi of trans-critorioyi chloride in 6 ml of lether was added chopwise, Alter stirring and cooling for 2 hours, the solvents were removed at reduced pressure. The residue was treated with valer and the resulting solid recrystalized from n-butanol to give 0,69 g of N-{4-{(3-bromophenyl)amino}-6-quinazolinyl-{6}-2-butenamide: mp = 183-160°C. Inass septemm (mrb. M+1 983, 984).

Example 8

N-[4-](3-Bromophenyl)amino]-6-quinazolinyl]-2-methyl-2-propenamide

A solution of 1.6 g of N-Q-bromophemy)-4.6-quickexelindiamine in 15 m fol pyritine was cooled in an ice bath and a solution of 0.59 m of methacrycyl chlorida in 6 m of either was accided dropwise. After stifring and cooling for 2 hours, the solvents were removed at reduced prossure. The restitute was treated with water and the resulting solid was taken up in n-butanof (warming). Addition of either to the cooled-solution gives 0.44 g of N-14-1(3-bromophenyl)amino)-6-quina-colinyl-2-emple-properation in p = 40-26°C, mass spectrum (rive). M-11-93°C, and the solid properties of t

Example 9

N-[4-f(3-Bromophenyl)amino]-6-quinazolinyl]-2-butynamide

A solution of 0.50 g of 2-bulynoic acid in 10 m of a tetrahydrofusan was cooled in an ice bath. A 0.79 ml portion of isobulyt-thio-forwants (allowine) by 0.65 ml portion of N-metryl-morpholine wise sided, After Bebott 1 minutes a solution of 1.6 g of N-(3-butmoc)-hemyl-4,5-quinazolindiamine in 10 mil of pyridice was added. The reaction was allowed to come to grow interpretary and size owneright. The sockwards were removed at reducing breast and the solid laws energystalface from n-bulanol (o give 1.07 g of N-[4-1(3-bromophary)]aminoj-8-quinazodinyl-2-bulynamido; maks spectrum (m/e): 31, 383.

Example 10

4-[/4-[(3-Bromophenyl)amino]-6-quinazolinyllamino]-4-oxo-(E)-2-butenoic acid

A 2.5 ml portion of 1 0 N aquebus socium hydroxido was added to 2.5 g of 4-[14-(G-broinopbinyl)saminof-equiriacipal planting-4-oxo-(G-2-buterior add stryls star (Example 5) in 25 m of shanol. After stirring for an hour, 2 1 ml of concentrated hydrochoics and was added, and the reaction was stirred an additional 2 hours. The resulting solid was recrystallized from houtage to 19.7 g of 4-[14-(G-Broinophenyl)aminoj-6-quiriazbinyl]eminoj-4-oxo-(E)-2-būterioic palcit mass spectrum (mgb). N+H 418.

Example 11

N-[4-[(3-Bromopheny])aminol-6-quinazolinyl]-2,4-hexadienamide

A solution of 0.67g of 2.4-hoxediency, axid in 10 ml of teirahydroturan was cooled in an iss bath. A 0.79 ml portion of isobutly inhoriormate followed by a 0.66 ml portion of N-melhyl morpholine were added. After about 1 militude a solution of 1.5 g of N-(2-bromopheny)-4.5-quinazolindiamine in 10 ml of pyridine was added. The reaction was allowed to come for prom temperature and sit rovernight. The solvents were removed at reduced pressure and the solid was recreasilized to give 1.0 g of N-4(2-bromopheny)siming-6-quinazoliny[2-broadeneander mg = 259-260°C.

Example 12

N-[4-[(3-Bromophenyl)amino]-6-quinazolinyl]-2-cyclopenteneamide

A solution of 0.43 g of 2-cyclopentenoic acid in 5ml of tetrahydrofuran vias cooled in an ice bath. A 0.49 ml portion of isoboxyl chiboroformate followed by a 0.41 ml portion of N-methyl morpholine were added. After about 1 minute a solution of 1.0 g of N-(3-bromophenyl)-4,6-quinacpinderaine in 10 ml of pyridine was added. The reaction was allowed.

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to come to room temperature and sir overnight. Another 0.5 equivalents of mixed anhydride was added. The mixture was attract of 5 hours. The solvents were moved at neduced pressure and the solid was purified by chromospharyly on silica gol to give 0.30 g of N-(4-((3-bromopharyl)paininto)-5-quinazolinyl)-2-cyclopenteneamide: mass spectrum (m/ 4/ 90 /4M-14).

Example 13

N-[4-[(3-Bromophenyl]amine]-6-quinazolinyl]-2-propenamide

A solution of 2.0 g of Nr3-broincphenyl)-4,6-quinazolindernine in 10 miol pyridine was cooled in an ice bath and a solution of 0.5 in oil acrypt phindrise in 30 miol either was added deposites at PC. After stirring a from temperature for 3.5 hours, the solvents were removed at reduced pressure. The residue was purified by chromatography to give 0.2 at 0.1 Nr4-(4-broincohlarohlarino)-6-duriazolinuth-2-properature in review. If will solve in the property of th

15 Example 14

N-[4-[(3-Bromophenyl)amino]-6-quinazolinyl]-(3-phenyl-2-propynamide)

A solution of 0,95 p of 3-phproyl-2-protynois actile in 10ml of feltallydrium was cooled in an ice bath. A 0,92 at portion of losology theirodromate followed by a 0,50 ml portion of Normethyl indepolition were added. After about 1 minute assolution of 1,0 g of N-(6-brenophenyl)-4,6-gulnezelindjenine in 7 ml of pyridine was added. The reaction at 0°C for 1 hr. This activent years remayed at reduced pressure and the solid was positioned by chromatography on silica get if give 0.01 g of N-(4-16)-brenophenyl/panino)-6-quinazolinyl-(3-phonyl-2-propynamide): mass spectrum (m/e): 443.2, 443

Example 15

6-amino-4-chloroquinazoline

A mixture consisting of 3.25 g of 4-chioxo-6-nitroquinazoline, 10.8 g of sodium hydrosuffile, and 0.3 g of the phase transfer catalyst (C₂H₁₂SyNCH₂*C) in 97 ml of tetrahydrofuser and 32 ml of water was altired appliely for 2 hours: mixture was distinged with either and the organic layer was septembed. The organic solution was wiseled with brine and then dried over magnesium sultate. The solution was wasted, with prime and then dried over magnesium sultate. The solution was passed through a small column of allicat get. The solvent was removed at 30°C at reduced pressure giving 6-amino-4-thioroquinazoline which is used in the next step without additional oblification.

Example 16

[4-chloro-6-quinazollnyl]-2-butynamide

A solition of 1.64 g of Zivitynoie act in 46 m of steralydofuran was cooled in an ice bath. A 2.34 ml perign of sicolarly chloroformate followed by a 4.15 ml perign of Nembrily morpholine were selded. Alter about 10 minutes, his was poured into a solution of 6-amino-4-chloroquinazoline in 46 ml tetralydofuran. This mixture was sitired at corn temperature for Zhous. The mixture was poured into a mixture of brine and saturated actions followed sold mix his change of sold mixture of the sold mixture of sold mixture of sold mixture of the sold mixture of sold mixtur

Example 17

N-14-[(3-Bromophenyl)amino]-6-quinazolinyl]-2-butynamide

A solution consisting of 1.76 g of 4-chloro-6-quinazolimyl) 2-bulynamide and 1.23 g of 3-bromo artilline was refluxed under an inert atmosphere in 23 ml of isopropanol for 40 minutes. The mixture was cooled to room temperature and 200 ml of ether was acked giving 0.4 g of N4-ft-(3-bromophenyl)amino-6-quinazolinyl)-2-bulynamide as the hydrochloride sall. Neutralizing with sodium bicarbonate solution, extracting with child scelate, removal of the colvent), and recystalization from 1-bulenol gives N4-ft-(3-bromophenyl)amino-6-quinazolinyl-2-bulynamide as the free base,

Example 18

Nº-(4-Amino-2-cyanophenyl)-N,N-dimethylformamidine

A solution of 5.0 g (27.5 mmol) of N°-(2-cyano-4-nitrophenyl)-N,N-dimethyllomamidine, 39.9 g (41.8 ml, 412.4 mmol) of cyclohexene, and 0.8 g of 10% PdC in 980 ml of melinand was refluxed for 4 hrs. The hot mixture was fillered through Cellia. Solvent was removed and the residue was recrystalized from chloroform-carbon tetrachicle giving 4.9 g (89%) of the title compound as a light gray crystalline solid, mass spectrum (m/e)*198.9 (M+H, electrospiay).

Example 19

N-[3-Cyano-4-[[(dimethylamino)methylene]amino]phenyl]-2-butynamide

To a solution of 2.01 is Q3.9 mmol) of 2-bulynois acid and 2.9 ml (22.9 mmol) sickuly inhinotomratis in 50 ml introdycolomna was stime at 0.70 curier introgen as 2.4.2 g (2.8.3 mil., 22.8 mmol) of N-methyl impribitie was acidad over 3 min. After stiming for 15 min., a solution of N-14-amino-2-cyanophrayly-N,N-dimethyl-formamidine and 1.6 g (1.75 m.), 1.5.9 mmol) of N-methyl impropherien is 25 ml listality/dirotan was acided over 4 min. The mittor was stimed over 4 min. The mittor was stimed 50 min., at 0°C and 30 min. at room temperature. The mittorien was distored with 70 ml of strip acetate and powered into a mixture of brine and statuted solution blue shortest. The original layer was chief dis(9.5Cs) and filtered filtering in a part of allice gdt. The solvent was removed and the residue was stirred with 50 ml of ether. The suspended solid was callanded to other 5.5 in 4.85% of a of 9.95% of an off-white solid mass spitierrum (river): 25.50 (MP4), electrospierry.

Example 20

N-f4-f(3-Bromophenyl)aminoil-6-quinazolinyll-2-butynamide

A solution of 3.0 g (11.8 mmol) of N-[3-cyano-4-[(directitytemion)methylane]arinfol) phenylly2-butynemidde and (2.12.8 mmol) of 3-bronou antiller in 18 mil circular date was refused gently with eletring under piregen for 1 hr 15 min. The mixture was cooled in an loo bath and a solid mass formed. The solid was calleded by filtration and washed with either-actionities 1:1 to give a yellow solid which was recrystalized from eithernol giving 2.51 g of N/4-[18-bronoblanytaminol-6-duilargicitivil-2-butynamider mass specifyrin (firely: 931, 933.)

Claims

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1. A compound of the formula

wherein:

X is phanyl optionally substituted with one or more substituents selected from the group consisting of balogen, allyly of 1-6 carbon datons, alloxy of 1-6 carbon atoms, alloxy of 1-6 carbon atoms, alloxy of 2-7 carbon atoms, carbonally of 2-7 carbon atoms, carbonally of 2-7 carbon atoms, carbonally of 2-7 carbon atoms, armon, and alkenoylaminio of 1-6 carbon atoms, Rand FI, are each, independently, hydrogen, halogen, allyl of 1-6 carbon atoms, alloxy of 1-5 carbon atoms, hydroxy, or trilliprometrity;

He is hydrogen, alkyl of 1-6 carbon atoms, alkoxy of 1-6 carbon atoms, hydroxy, trifluoromethyl;

Y is a radical selected from the following:

R₃ is independently hydrogen, alkyl of 1-6 carbon atoms, carboxy, carboalkoxy of 1-6 carbon atoms, phenyl, or carboalkyl of 2-7 carbon atoms:

n = 2-4; or a charmaceutically acceptable saft thereof, with the proviso that each R_1 of Y may be the same or different.

- 2. Acompound according to Claim 1 wherein R, R₁, and R₂ are hydrogen or a pharmaceutically acceptable salt thereof
- 3. A compound according to Claim 1 or Claim 2 wherein X is unsubstituted or substituted with halogen or alkyl of 1-6. carbon atoms.
- 4. A compound according to Claim 1 which is one of the following :

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- N-[4-[(3-Bromophenyl)amino]-6-quinazolinyl]-2-butynamide;
- N-[4-[(3-Bromophenyl)amino]-6-quinazolinyli-2-methyl-2-propenamide;
- N-(4- [(3-Bromophenyl)amino]-6-quinazolinyl]-2,4-hexadienamide;
- N-[4-[(3-Bromophenyl)amino]-6-quinazolinyl]-(E)-2-butenamide;
 - N-[4-[(3-Bromophenyl)amino]-6-quinazolinyl]-3-methyl-2-butenamide;
 - 4-[[4-[(3-Bromophenyl]amino]-6-quinazolinyl]amino]-4-oxo-(Z)-2-butenoic acid; 4-[[4-[(3-Bromophenyl)amino]-6-quinezolinyl]amino]-4-oxo-(E)-2-butenoic acid;
- 4-[[4-[(3-Bromophenyl]amino]-6-quinazolinyl]amino]-4-oxo-(E)-2-butencic acid, ethyl ester;
- N-[4-[(3-bromophenyl)amino]-6-quinazolinyl]-2-cyclopentenamide;
- N-[4-[(3-Bromophenyl)amino]-6-quinazolinyl]-2-propenamide

- N-[4-((3-bromophenyl)amino]-6-quinazolinyl]-(3-phenyl-2-propynamide); or a pharmaceutically acceptable salt thereof.
- A method of inhibiting the biological effects of a deregulated protein tyrosine kinase in a mammal which comprises administering to said mammal an effective amount of a compound having the formula 1 as defined in claim 1 or a pharmaceutically acceptable salt thereof.
- 6. A method of treating, inhibiting the growth of, or eradicating neoplasms in a mammal which comprises administering to said mammalan effective amount of a compound having the formula 1 as defined in claim 1 or a pharmaceutically acceptable salt thereof.
- A method according to claim 6 wherein the necolasm expresses EGFR.
 - 8. A method according to claim 7 wherein the neoplasm is selected from the group consisting of breast, kidney, bladder, mouth, larvnx, esophagus, stomach, colon, ovary, and lung.

- A pharmaceutical composition which comprises a compound having the formula 1 as defined in claim 1 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.
- 10. A process for preparing a compound of the formula:

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wherein R, R, R, and X are as defined in claim 1; Y is a radical selected from the following:

$$R_3$$
 R_3 R_3 R_3 R_4 R_5 R_6 R_8 R_8

$$R_3$$
 R_3 R_3 R_3 R_3 R_3 R_3 R_3

wherein each H₃ is independently hydrogen, alkyl of 1-6 carbon atoms, carboxy, carboxyl of 2-6 carbon atoms, phenyl, or carboalkoxy of 2-7 carbon atoms; it is the integer 2-4, which comprises one of the following:

a) reacting a 6-amino-quinazoline of the formula:

wherein X, R, R₁ and R₂ are as defined above, with an acid chloride or mixed anhydride of the formula:

wherein Y is as defined above and P4 is alkyl of 1-6 carbon atoms to give a compound of formula 1;

b) reacting a compound of formula 6 as defined above with a compound of formula 11

wherein each H₈ is independently hydrogen, phenyl or alkyl of 1 to 6 carbon aforms, to give a compound as claimed in claim 1 having the formula 12:

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wherein X, H, H, H, H2 and H5 are as defined above;

c) reacting a compound of the formula:

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wherein Y is as defined above, with an aniline of the formula:

wherein \boldsymbol{X} is as defined in Claim 1 to give a compound of formula 1; or

d) reacting a compound of the formula:

wherein Y is as defined above, with an aniline of the formula:

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X-NH_{2 18}

wherein X is as defined in Claim 1, to give a compound of formula 1; and, if desired after any of steps a), b), c) and d), isolating the product as a pharmaceutically acceptable salt.

11. A process as claimed in claim 10 in which the compound of formula

as defined in Claim 10 is prepared by reducing a compound of formula

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wherein R, R, R, R, and X are as defined in Claim 1.

12. A process for as claimed in Claim 11 in which the compound of the formula:

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wherein X, R, R₁, and R₂ are as defined in Claim 1, is prepared by reacting a compound of formula:

$$O_2N$$
 O_2N
 O_2N

with an aniline of the formula:

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X-NH₂₄

wherein R, R, R, and X are as defined in Claim 11,

6 13. A process for preparing a compound of the formula 1 as defined in Claim 1 which comprises reacting an anthranilonitrile of the formula:

with dimethylformamide dimethyl acetal to give a compound of the formula:

heating said compound with an aniline of the formula:

X-NH

in an acidic organic solvent to give a 6-nitro-quinazoline of the formula:

5,

treating said compound with a reducing agent to give a 6-amino-quinazoline of the formula:

and reacting said compound with an acid chloride or mixed anhydride of the formulas:

wherein R₄ is alkyl of 1-5 carbon atoms, and if desired isolating as a pharmaceutical sait.

14. A process for producing a compound of the formula:

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wherein X and Y are according to claim 1; which comprises reducing a compound of the formula;

with sodium hydrosulfite and a phase transfer cafetyst in a solvent mixture comprising an inert organic solvent and water to give a compound of the formula:

reacting said compound with an acid chloride or mixed anhydride of the formula:

wherein Y is as defined in Claim 1 and wherein H_a is alkyl of 1-5 carbon atoms, to give a compound of the formula:

and reacting said compound with an aniline of the formula:

X-NH.

15. A precess for producing a compound of the formula:

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wherein X and Y are as according to claim 1; which comprises reducing the compound of the formula:

with a palladium catalyst and a source of hydrogen in an inert solvent to give a compound of the formula:

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reacting said compound with an acid chloride or mixed anhydride of the formula:

wherein H₄ is alkyl of 1-6 carbon atoms, to give a compound of the formula:

and reacting said compound with an aniline of the formula:

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X-NH₂



PARTIAL EUROPEAN SEARCH REPORT Application Number

which under Rule 45 of the European Patent Convention EP 97 30 0703 shall be considered, for the purposes of subsequent proceedings, as the European Search report

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